What is claimed is:

- 1. A method comprising the steps of: (a) providing a sample suspension containing at least one set of G protein beads, each of said G-protein beads comprising epitoperecognizing beads having a heterotrimeric G protein bound thereto; (b) mixing said sample suspension with at least one type of G protein coupled receptor and a ligand to thereby form a mixed suspension containing a ligand-receptor-G protein complex when said G protein is capable of forming a complex with said receptor and said ligand; (c) mixing said mixed suspension to incubate said mixed suspension and thereby form an incubated suspension; (d) detecting the formation of a stable ligand-receptor-G protein complexes on the said beads in said incubated suspension by flow cytometry.
- 2. The method of claim 1, further comprising the step of determining whether said G protein is capable of forming a complex with said receptor and said ligand, wherein said receptor and said ligand are known to form a complex with an unknown G protein.
- 3. The method of claim 1, further comprising the step of determining whether said ligand is capable of forming a complex with said G protein and said receptor, wherein said G protein and said receptor are known to form a complex with another ligand.
- 4. The method of claim 1, further comprising forming said G protein beads by binding epitope-bearing G protein subunits to epitope-recognizing beads.
- 5. The method of claim 1, wherein said eptitope-recognizing beads comprise beads that recognize the FLAG epitope.
- 6. The method of claim 1, wherein said epitope-recognizing beads comprise beads that recognize the six-histidine (H6) epitope.
- 7. The method of claim 1, further comprising the step of solubilizing said receptor.

- 8. The method of claim 1, wherein said epitope-recognizing beads are derivatized to carry chelated nickel.
- 9. The method of claim 1, wherein the beads displaying said heterotrimeric G protein include a fluorescent address tag.
- 10. The method of claim 1, wherein said ligand includes a fluorescent tag.
- 11. The method of claim 1, wherein said receptor is fluorescent.
- 12. The method of claim 11, wherein said receptor is made fluorescent by chemical derivatization of said receptor.
- 13. The method of claim 11, wherein said receptor is made fluorescent by making a fusion protein of said receptor with a fluorescent protein.
- 14. The method of claim 1, wherein said at least one set of G protein beads comprises at least two sets of G protein beads, each of said sets of G protein beads having a different respective fluorescent color and a different respective heterotrimeric G protein bound thereto; wherein all of said sets of G protein beads are simultaneously present in said mixed suspension; and wherein step (d) comprises detecting the respective fluorescent color associated with each set of G protein beads separately.
- 15. The method of claim 14, wherein said at least one type of G protein coupled receptor comprises at least two types of G protein coupled receptors.
- 16. The method of claim 1, wherein said at least one type of G protein coupled receptor comprises at least two types of G protein coupled receptors, each of said types of G protein coupled receptors having a different fluorescent color; wherein all of said types of G protein coupled receptors are simultaneously present in said mixed suspension; and wherein step (d) comprises detecting the respective fluorescent color associated with each

of said types of G protein coupled receptors separately.

- 17. The method of claim 16, wherein G protein beads have a first fluorescent color, wherein said ligand is bound to ligand bearing beads having a second fluorescent color, wherein said G protein beads and said ligand bearing beads are present simultaneously in said mixed suspension, and wherein step (d) comprises detecting the respective fluorescent color associated with said G protein beads and said ligand bearing beads separately.
- 18. The method of claim 1, wherein a regulator of G protein signaling is mixed with said mixed suspension and wherein said method further comprises the following step: (e) determining the effect of said regulator of G protein signaling upon ternary complex formation.
- 19. The method of claim 18, wherein said at least one set of G protein beads comprises at least two sets of G protein beads, each of said sets of G protein beads having a different respective fluorescent color and a different respective heterotrimeric G protein bound thereto; wherein all of said sets of G protein beads are simultaneously present in said mixed suspension; wherein step (d) comprises detecting the respective fluorescent color associated with each set of G protein beads separately, and wherein step (e) comprises determining the effect of said regulator of G protein signaling upon ternary complex formation for each of said sets of G protein beads.
- 20. The method of claim 19, wherein said at least one type of G protein coupled receptor comprises at least two types of G protein coupled receptors and wherein step (e) comprises determining the effect of said regulator of G protein signaling upon ternary complex formation for each of said types of G protein coupled receptors.
- 21. The method of claim 18, wherein said at least one type of G protein coupled receptor comprises at least two types of G protein coupled receptors, each of said types of G protein coupled receptors having a different fluorescent color; wherein all of said types of

G protein coupled receptors are simultaneously present in said mixed suspension; wherein step (d) comprises detecting the respective fluorescent color associated with each of said types of G protein coupled receptors separately; and wherein step (e) comprises determining the effect of said regulator of G protein signaling upon ternary complex formation for each of said types of G protein coupled receptors.

- 22. A method claim 18, comprising forming said G protein beads by the steps of: providing epitope-recognizing beads; and binding epitope-bearing G protein subunits to said epitope-recognizing beads to form G protein beads.
- 23. The method of claim 22, wherein said eptitope-recognizing beads comprise beads that recognize the FLAG eptitope.
- 24. The method of claim 22, wherein said eptitope-recognizing beads comprise beads that recognize the six-histidine (H6) epitope.
- 25. The method of claim 22, wherein said epitope-recognizing beads are derivatized to canny chelated nickel.
- 26. The method of claim 22, wherein said heterotrimeric G protein includes a fluorescent tag.
- 27. A method comprising evaluating a G protein coupled receptor agonism or antagonism of a compound by a bead-based flow cytometric process comprising contacting the compound with beads conjugated to a G protein coupled receptor ligand which would result in a detectable G protein coupled receptor ligand-receptor complex, to determine the existence of an interaction or an an absence of an interaction with said receptor, wherein the extent to which the compound competetes with said ligand-receptor complex determines that a compound is an agonist or an antagonist of said G protein coupled receptor.

- 28. A method comprising evaluating a β 2-adrenergic receptor agonism or antagonism of a compound by a bead-based flow cytometric process comprising contacting the compound with beads conjugated to a β 2-adrenergic receptor ligand which would result in a detectable β 2-adrenergic receptor ligand-receptor complex, to determine the existence of an interaction or an an absence of an interaction with said receptor, wherein the extent to which the compound competes with said ligand-receptor complex determines that a compound is an agonist or an antagonist of said β 2-adrenergic receptor.
- 29. A method of claim 28, wherein the beads are dihydroalprenolol-conjugated beads.
- 30. A method comprising evaluating a G-protein receptor agonism or partial agonism of a compound in a bead based high throughput screening system comprising a) contacting the compound and solubilized detectable G protein coupled receptor with G protein beads, each of said G-protein beads comprising epitope-recognizing beads having an epitope-tagged heterotrimeric G protein bound thereto; and b) determining whether a ternary complex between said G protein coupled receptor and said G protein occurs, wherein an interaction between receptor and G protein evidences that said compound is an agonist or partial agonist of said G protein coupled receptor.
- 31. A method of claim 30, wherein the G protein receptor is the β 2-adrenergic receptor, said receptor contains a fluorescent moiety, and the interaction between said receptor and said G protein evidences that said compound is an agonist of said receptor.
- 32. A method of claim 30, wherein said the detectable moiety is a any fluorescent protein.
- 33. A method of claim 31, wherein the β 2-adrenergic receptor is a β 2AR-GFP fusion protein.

- 34. A method of claim 30, wherein detectable ternary complex levels are used to generate dose-response curves that are indicative of the compound's β 2-adrenergic receptor agonism or antagonism.
- 35. A method of claim 30, wherein GTP γ S-induced activation rates for the detectable ternary complex are determined and wherein compounds that are β 2-adrenergic receptor agonists or partial β 2-adrenergic receptor agonists have approximately equal GTP γ S-induced activation rates.
- 36. A method comprising evaluating the relative G protein receptor agonism or partial agonism a compound by a flow cytometric process comprising contacting the compound and soluble detectable G protein receptor with beads conjugated to epitope-recognizing beads having a heterotrimeric G protein bound thereto, wherein an agonist or partial agonist compound binds to G protein receptor to form a compound-receptor complex and said compound-receptor complex binds to said bound G protein to form a detectable ternary complex indicative of the compound's G protein receptor agonism or antagonism.
- 37. A method of claim 36, wherein the G protein receptor is a β 2-adrenergic receptor containing a fluorescent moiety.
- 38. A method of claim 37, wherein the fluorescent moiety is a GFP or a RFP fused to said G protein receptor.
- 39. A method of claim 36, wherein the detectable β 2-adrenergic receptor is a β 2AR-GFP fusion protein.
- 40. A method of claim 36, wherein GTP γ S-induced activation rates for the detectable ternary complex are determined and wherein compounds that are β 2-adrenergic receptor agonists or partial β 2-adrenergic receptor agonists have approximately equal GTP γ S-induced activation rates.

- 41. A method of evaluating a library of compounds comprising: selecting a plurality of compounds from the library; evaluating the relative $\beta 2$ -adrenergic receptor agonism of each selected compound by a flow cytometric process comprising contacting the compound with beads conjugated to a $\beta 2$ -adrenergic receptor-detectable moiety complex, wherein the extent to which the compound complexes with the $\beta 2$ -adrenergic receptor-detectable moiety complex to form a detectable ternary complex is determined by measuring detectable ternary complex levels and detectable ternary complex levels are indicative of the compound's $\beta 2$ -adrenergic receptor agonism or antagonism; and evaluating a differentiation state or a metabolic parameter of the cell or organism.
- 42. A method comprising evaluating the relative G protein receptor agonism, antagonism or inactivity of a compound for a G protein coupled receptor (GPCR) in a single sample by a flow cytometric process comprising the steps of (a) providing a sample suspension containing a detectable GPCR, a set of G protein beads which will form a ternary complex with said detectable GPCR in the presence of an agonist or partial agonist, and a set of ligand beads which will bind to said detectable GPCR, said set of G-protein beads comprising epitope-recognizing beads having a heterotrimeric G protein bound thereto; (b) mixing said sample suspension with said compound; and (c) detecting the formation or absence of formation of a complex between said compound and said detectable GPCR, wherein a GPCR antagonist prevents binding of said detectable GPCR to said G protein beads by preventing ternary complex formation and prevents binding of said detectable GPCR to said ligand bead; a GPCR agonist allows binding of said detectable GPCR to said G protein beads by forming a ternary complex but prevents binding of said detectable GPCR to said ligand bead; and an inactive compound prevents binding of said detectable GPCR to said G protein beads by not promoting ternary complex formation but allows binding of said detectable GPCR to said ligand bead.
- 43. A method of claim 42, wherein the G protein coupled receptor is a β 2-adrenergic receptor containing a fluorescent moiety.

- 44. A method of claim 42, wherein the fluorescent moiety is any fluorescent protein fused to said G protein coupled receptor.
- 45. A method of claim 43, wherein the detectable β 2-adrenergic receptor is a β 2AR-GFP fusion protein.
- 46. The method of claim 42 wherein said G protein beads are modified with a fluorescent moiety.
- 47. The method of claim 46 wherein said fluorescent moiety is Texas Red.
- 48. A method comprising identifying agents useful in the treatment of a disease associated with G protein coupled receptor (GPCR) agonism or antagonism by determining an agent's GPCR agonism or antagonism by a flow cytometric process comprising: (a) providing a sample suspension containing a detectable GPCR, a set of G protein beads which will form a ternary complex with said detectable GPCR in the presence of an agonist or partial agonist, and a set of ligand beads which will bind to said detectable GPCR, said set of G-protein beads comprising epitope-recognizing beads having a heterotrimeric G protein bound thereto; (b) mixing said sample suspension with said agent; and (c) detecting the formation or absence of formation of a complex between said agent and said detectable GPCR, wherein a GPCR antagonist prevents binding of said detectable GPCR to said ligand bead; and a GPCR agonist allows binding of said detectable GPCR to said G protein beads by forming a ternary complex but prevents binding of said detectable GPCR to said ligand bead.
- 49. A method of claim 48, wherein the G protein receptor is a β 2-adrenergic receptor containing a fluorescent moiety.
- 50. A method of claim 48, wherein the fluorescent moiety is any fluorescent protein fused to said G protein receptor.

- 51. A method of claim 49, wherein the detectable β 2-adrenergic receptor is a β 2AR-GFP fusion protein.
- 52. The method of claim 48 wherein said G protein beads are modified with a fluorescent moiety.
- 53. The method of claim 46 wherein said fluorescent moiety is Texas Red.
- 54. A method comprising evaluating a G protein coupled receptor agonism or antagonism of a compound by a bead-based flow cytometric process comprising contacting the compound with beads conjugated to a ligand which would result in a detectable G protein coupled receptor ligand-receptor complex, to determine the existence of an interaction or an an absence of an interaction with said detectable receptor, and comparing said interaction with said ligand-receptor complex with a ligand-receptor complex utilizing a known agonist or antagonist to determine that a compound is an agonist or an antagonist of said G protein coupled receptor.
- 55. A method of evaluating a library of compounds comprising: selecting a plurality of compounds from the library; evaluating the relative $\beta 2$ -adrenergic receptor agonism or antagonism of each selected compound by a flow cytometric process comprising contacting the compound with beads conjugated to a ligand for a $\beta 2$ -adrenergic receptor-detectable moiety complex, wherein the extent to which the compound complexes with the $\beta 2$ -adrenergic receptor-detectable moiety and binds to said ligand conjugated beads is indicative of agonist or antagonist activity; and evaluating a differentiation state or a metabolic parameter of the cell or organism.